

METHOD AND DEVICE USING MYOELECTRICAL ACTIVITY FOR  
OPTIMIZING A PATIENT'S VENTILATORY ASSIST

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FIELD OF THE INVENTION

The present invention relates to a method and device for determining a level of ventilatory assist to a ventilator-dependent patient.

BACKGROUND OF THE INVENTION

10 Both the tension developed by a patient's muscle [34] and the duration of the muscle contraction [2] are factors that lead to respiratory muscle fatigue; these two factors can be expressed by indices such as the tension-time index [3] and the pressure-time product [10, 20, 32, 35]. Bellemare and Grassino [3] showed a direct inverse relationship exists between the time of endurance of a  
15 fatiguing diaphragm contraction and the rate of decay of the ratio of the high to low spectral components (H/L) of the electrical activity EAdi of the patient's diaphragm, indicating that these two values are indicative of progressive failure to sustain load. The force exerted by the muscle has been shown to be directly related to the rate of decay of the power spectrum center frequency or the rate  
20 of decay of the above mentioned ratio H/L, and the level at which this power spectrum center frequency or ratio H/L plateaus [16, 21, 28]. Such shifts in the power spectrum reflect a reduction in the muscle action potential conduction velocity [28, 38, 39], and constitute an early indication that, at the cellular level, these breathing patterns cannot be maintained indefinitely [3].

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Hyperinflation, which impairs the length-tension relationship of the respiratory muscles, i.e. the transformation of the neural activation into a mechanical output or pressure, reduces the capacity of the respiratory muscles to generate pressure (neuromechanical uncoupling), unless the electrical activity  
30 EAdi of the patient's diaphragm is increased. Studies have shown that when the

inspiratory pressure, flow and duty cycle remain constant, increases in end-expiratory lung volume (EELV) promote reductions in endurance time [33, 44] and sustainable pressure [11]. In an animal model, Tzelepis et al [44] proposed that, under these conditions, diaphragm shortening would require greater 5 excitation to generate a given sub-maximum tension, and that this increased excitation might account for the greater contractile muscle fatigability observed at shorter muscle length.

10 The level of partial ventilatory assist, with the aim to ensure adequate pulmonary ventilation while preserving inspiratory muscle function, is generally set on an empirical basis in the clinical setting.

15 It has been proposed that an optimal level of partial ventilatory assist could be determined from the lowest stable breathing frequency  $f_B$  achieved, i.e. without bradypnea or apnea. In patients, this corresponded to 16.4 bpm (breaths per minute) and was associated with a tidal volume  $V_T$  of 11.8 ml/kg. However, mechanical lung modeling in that study demonstrated that such a level of support actually resulted in a total unloading of the respiratory muscles.

20 Others have defined an optimal level of partial ventilatory assist as that which produces the lowest swings of transdiaphragmatic pressure  $P_{di}$  and found that this condition was associated with a breathing frequency  $f_B$  of 19.7 bpm and a tidal volume  $V_T$  of 11.7 ml/kg. The transdiaphragmatic pressure  $P_{di}$  in the latter study was used as a marker of inspiratory effort.

25 Jubran et al [20] defined an upper bound inspiratory pressure-time product lower than 125 cm H<sub>2</sub>O·s/min as a desirable level of inspiratory effort to be achieved during partial ventilatory assist. Although arbitrarily determined, this threshold was justified by the fact that it corresponded to a tension-time index 30  $TT_{di}$  well below that considered to indicate impeding inspiratory muscle fatigue. The study found a high variability in pressure-time products between patients and demonstrated that a breathing frequency  $f_B < 30$  bpm and a tidal volume  $V_T$

of 0.6 L were better determinants of an optimal level of inspiratory effort during partial ventilatory assist. Based on these breathing pattern findings, it is likely that the level of respiratory muscle unloading provided by this method of optimizing partial ventilatory assist was lower than that of the above discussed  
5 studies.

Brochard et al [8] defined an optimal partial ventilatory assist level as the lowest level of ventilatory assist, which when implemented, maintained the highest level of diaphragmatic electrical activation without the occurrence of  
10 fatigue as evaluated via power spectrum analysis of the electrical activity  $EA_{di}$  of the patient's diaphragm. Interestingly, such levels of partial ventilatory assist were associated with a breathing frequency  $f_B$  of 20-27 bpm and a tidal volume  $V_T$  of 8.0 ml/kg, these values being similar to those later reported by Jubran et al [20].  
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#### SUMMARY OF THE INVENTION

In accordance with the present invention, there is provided a method for  
20 determining a level of ventilatory assist to a ventilator-dependent patient, comprising: calculating a critical threshold of a respiration-related feature, wherein fatigue of a respiratory muscle of the ventilator-dependent patient develops when the critical threshold is reached by the respiration-related feature; and controlling the level of ventilatory assist to the ventilator-dependent  
25 patient in relation to the critical threshold of the respiration-related feature so as to prevent fatigue of the patient's respiratory muscle.

The present invention also relates to a device for determining a level of ventilatory assist to a ventilator-dependent patient, comprising: a calculator of a  
30 critical threshold of a respiration-related feature, wherein fatigue of a respiratory muscle of the ventilator-dependent patient develops when the critical threshold is reached by the respiration-related feature; and a controller of the level of

ventilatory assist to the ventilator-dependent patient in relation to the critical threshold of the respiration-related feature so as to prevent fatigue of the patient's respiratory muscle.

5       The foregoing and other objects, advantages and features of the present invention will become more apparent upon reading of the following non-restrictive description of illustrative embodiments thereof, given by way of example only with reference to the accompanying drawings.

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#### BRIEF DESCRIPTION OF THE DRAWINGS

In the appended drawings:

15       Figure 1 is a schematic representation of a non-limitative example of experimental set-up for measuring diaphragm's electrical activity EAdi, esophageal pressure  $P_{es}$ , gastric pressure  $P_{ga}$ , respiratory airflow and tidal volume  $V_T$ , and for displaying on line the target transdiaphragmatic pressure  $P_{di}$  and the root-mean-square (RMS) of the diaphragm's electrical activity EAdi;

20       Figure 2 are illustrative examples of tracings of tidal volume  $V_T$ , diaphragm electrical activity EAdi, transdiaphragmatic pressure  $P_{di}$ , esophageal pressure  $P_{es}$ , and gastric pressure ( $P_{ga}$ ) measured on a subject during "volume" maneuvers and "expulsive" maneuvers;

25       Figure 3 are examples of bar graphs displaying drops in center frequency  $CF_{di}$ , targeted levels of transdiaphragmatic pressure  $P_{di}$ , diaphragm pressure-time product  $PTP_{di}$  and the associated diaphragm's electrical activity EAdi observed during volume and lower-pressure expulsive and higher pressure expulsive maneuvers;

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      Figure 4 are examples of graphs from one representative subject showing the center frequency  $CF_{di}$ , the root-means-square (RMS) of the

diaphragm's electrical activity EAdi and the diaphragm pressure-time product PTP<sub>di</sub> plotted over time during the volume maneuver (circles) and the two expulsive maneuvers at end-expiratory lung volume (EELV), one targeting a lower transdiaphragmatic pressure P<sub>di</sub> (squares) and the other a higher P<sub>di</sub>

5 transdiaphragmatic pressure P<sub>di</sub> (triangles);

Figure 5 is a flow chart and block diagram of a first non-restrictive illustrative embodiment of the method and device according to the present invention, for determining a level of ventilatory assist to a ventilator-dependent  
10 patient; and

Figure 6 is a flow chart and block diagram of a second non-restrictive illustrative embodiment of the method and device according to the present invention, for determining a level of ventilatory assist to a ventilator-dependent  
15 patient.

#### DETAILED DESCRIPTION OF THE ILLUSTRATIVE EMBODIMENTS

A study was conducted to determine in humans whether an increased  
20 electrical activity EAdi of a patient's diaphragm, with neuromechanical uncoupling, promotes greater reductions in the center frequency CF<sub>di</sub> of the diaphragm's electrical activity EAdi, when the diaphragm pressure-time product PTP<sub>di</sub> is kept constant. An additional aim of the study was to establish the extent to which the diaphragm pressure-time product PTP<sub>di</sub> needs to be increased, in  
25 the presence of normal neuromechanical coupling, in order to reproduce the drop in center frequency CF<sub>di</sub> observed with uncoupling.

More specifically, the study evaluated whether increased diaphragm activation induced by an increased lung volume promotes increased drops in the  
30 center frequency CF<sub>di</sub> of the diaphragm's electrical activity EAdi when the diaphragm pressure-time product PTP<sub>di</sub> is kept constant. Five healthy subjects performed runs of intermittent quasi-static diaphragmatic contractions with a

fixed breathing pattern. In separate runs, the subjects targeted transdiaphragmatic pressures  $P_{di}$  by performing end-inspiratory holds at total lung capacity with the glottis open (neuromechanical uncoupling), and at end-expiratory lung volume by performing expulsive maneuvers (no neuromechanical uncoupling). Diaphragm activation and pressures were measured with an electrode array and with balloons, respectively, mounted on an esophageal catheter. Reproduction of a transdiaphragmatic pressure  $P_{di}$  of  $\approx 31$  cm H<sub>2</sub>O during neuromechanical uncoupling increased lung volume to 77.5% of the inspiratory capacity, increased the diaphragm's electrical activity EAdi from 25% to 61% of the maximum and resulted in a 17% greater drop in center frequency CF<sub>di</sub>. In order to reproduce, in the absence of neuromechanical uncoupling, the decrease in center frequency CF<sub>di</sub> observed during neuromechanical uncoupling, a two-fold increase in transdiaphragmatic pressure P<sub>di</sub> and diaphragm pressure-time product PTP<sub>di</sub> was required. It was concluded that a constant diaphragm pressure-time product PTP<sub>di</sub> does not necessarily result in a center frequency CF<sub>di</sub> of the diaphragm's electrical activity EAdi that remains stable when activation is increased.

## METHODS

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### *Subjects*

Five healthy subjects (1 female, 4 males) with a mean age of  $40.6 \pm 8.0$  years participated in the study. The study was approved by the Scientific and Ethical Committees of Sainte-Justine's Hospital and all subjects gave their informed consent.

### *Experimental protocol*

30 Figure 1 is a schematic representation of a non-limitative example of experimental set-up. On the left, an esophageal catheter-mounted multi-electrode array 5 is used to measure diaphragm's electrical activity EAdi and

balloons 8 and 9 mounted on the catheter on opposite sides of the electrode array 5 are used to measure esophageal pressure  $P_{es}$  and gastric pressure  $P_{ga}$ . The catheter 6 was passed trans-nasally and positioned at the gastro-esophageal junction 10. Respiratory airflow was measured with a 5 pneumotachograph 3 and tidal volume  $V_T$  was obtained by integrating inspiratory flow. On the right, the target transdiaphragmatic pressure  $P_{di}$  and the root-mean-square (RMS) of the diaphragm's electrical activity EAdi are displayed on line.

Referring to Figure 1, each human subject 2, while seated in an upright 10 chair (not shown) and facing a the monitor 1 of a computer 4, performed repeated maximal inspirations to total lung capacity (TLC) in order to obtain three reproducible voluntary maximum values for the diaphragm's electrical activity EAdi. Each subject 2 was subsequently asked to perform intermittent, near-isometric diaphragmatic contractions of 10 seconds duration, separated by 15 5 seconds relaxation periods during which free breathing was allowed. With visual feedback of the transdiaphragmatic pressure  $P_{di}$  on the monitor 1 of the computer 4 a low level of transdiaphragmatic pressure  $P_{di}$  was targeted during two runs, while a higher level of transdiaphragmatic pressure  $P_{di}$  was targeted during a third run. The duty cycle was imposed by a sound signal, and each run 20 lasted until a plateau in center frequency  $CF_{di}$  was reached, or until the subject was no longer able to maintain the target transdiaphragmatic pressure  $P_{di}$ .

Figure 2 are examples of tracings of tidal volume  $V_T$ , diaphragm electrical 25 activity EAdi, transdiaphragmatic pressure  $P_{di}$ , esophageal pressure  $P_{es}$ , and gastric pressure  $P_{ga}$  measured in one subject during "volume" maneuvers and "expulsive" maneuvers performed during the hereinafter reported study. The "volume" maneuver consisted of an end-inspiratory hold at an increased lung volume, which resulted in the generation of a low  $P_{di}$  (left tracing), whereas the two expulsive maneuvers were performed at end-expiratory lung volume 30 targeting a lower  $P_{di}$  (middle tracing) and higher  $P_{di}$  (right tracing).

In order to obtain two different levels of diaphragm's electrical activity

EAdi for the same target transdiaphragmatic pressure  $P_{di}$ , each subject 2 was instructed to perform two different maneuvers:

1. Volume maneuver: the subjects inspired close to their total lung capacity (TLC) and produced a given level of transdiaphragmatic pressure  $P_{di}$  (Figure 2; left tracing). The transdiaphragmatic pressure  $P_{di}$  was maintained at this lung volume with the glottis open.
  2. Expulsive maneuver: the subjects performed expulsive maneuvers in order to generate a target transdiaphragmatic pressure  $P_{di}$ . All expulsive maneuvers were performed at end-expiratory lung volume (EELV) at lower and higher transdiaphragmatic pressures  $P_{di}$  (Figure 2, middle and right tracings).
- 15        After having initially performed a volume maneuver run, each subject 2 then performed two expulsive maneuver runs. One expulsive maneuver run targeted a transdiaphragmatic pressure  $P_{di}$  (lower pressure) similar to that observed during the volume maneuver but requiring less diaphragm's electrical activity EAdi, while another expulsive maneuver run targeted an increased 20 transdiaphragmatic pressure  $P_{di}$  (higher pressure) to reproduce the center frequency  $CF_{di}$  observed during the volume maneuver run. The volume maneuver was subsequently repeated once for retest purpose. The subject rested for 20 minutes between subsequent runs.

25        *Instrumentation*

Using the set-up of Figure 1:

- airflow and tidal volume were measured by a computer 4 through a 30 pneumotachograph 3;
- electrical activity EAdi of the patient's diaphragm was measured by the

computer 4 through the linear array 5 of electrodes mounted on an esophageal catheter 6 inserted through the patient's nostril (or patient's mouth) until the electrode array 5 is positioned in the gastro-esophageal junction 10 of the patient's diaphragm 7;

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- esophageal  $P_{es}$  and gastric  $P_{ga}$  pressures were measured by the computer 4 through the gastric 8 and esophageal 9 balloons mounted on the catheter 6 on opposite sides of the array 5 of electrodes; and
- 10 - the transdiaphragmatic pressure  $P_{di}$  was obtained by the computer 4 by subtracting the measured esophageal pressure  $P_{es}$  from the measured gastric pressure  $P_{ga}$ .

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*On-line automatic processing of diaphragm's electrical activity EAdi*

The diaphragm's electrical activity EAdi, more specifically a root-mean-square (RMS) EAdi signal was acquired, processed and displayed on-line using a standardized methodology [4, 36, 41]. The center frequency  $CF_{di}$  was evaluated for signal quality using established indices and criteria in accordance with a method disclosed by Sinderby et al [40]. To avoid influence of power spectral shifts on the EAdi signal strength, the RMS EAdi signal was calculated on the spectral moment of order 1 (M1) which is insensitive to conduction velocity [6] (see upper trace on the computer monitor 1. For more extensive review reference is made to Aldrich et al [1].

*Off-line signal analysis*

Inspiratory duration  $T_i$ , total breath duration  $T_{tot}$ , and breathing frequency 30  $f_B$ , diaphragm's electrical activity EAdi and pressures  $P_{es}$  and  $P_{ga}$  were determined using the transdiaphragmatic pressure  $P_{di}$ . The diaphragm pressure-time product  $PTP_{di}$  was obtained by multiplying (i) the under-the-curve area

subtended by the  $P_{di}$  signal by (ii) the breathing frequency  $f_B$ . The amplitude of the signal of the diaphragm's electrical activity  $EAdi$  was expressed as a percentage of the voluntary maximum diaphragm's electrical activity  $EAdi$  obtained from TLC maneuvers [37]. Variables were compared between each of 5 the maneuvers performed using one-way repeated measurements analysis of variance (ANOVA) and post hoc contrasts of significant effects were performed using the Student-Newman-Keuls test. Test-retest reliability of the  $P_{di}$ ,  $EAdi$  and  $CF_{di}$  values obtained during the volume and expulsive maneuvers was evaluated by calculating the interclass correlation coefficient (ICC).

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## RESULTS

The subjects were able to perform all maneuvers and maintain the imposed duty cycle ( $P=0.93$ ; one -way ANOVA) during all protocols (Table 1).

15

Table 1

*Breathing pattern and targeted  $P_{di}$  values during the three maneuvers performed*

Subject	Volume maneuver				Expulsive maneuver		Expulsive maneuver	
					Lower pressure		Higher pressure	
	Ti/Ttot	Vt%IC	Pdi	Ti/Ttot	Pdi	Ti/Ttot	Pdi	
1	0.66	89.2	47.8	0.66	45.1	0.67	77.3	
2	0.66	65.6	25.3	0.67	26.6	0.66	93.3	
3	0.67	68.3	10.2	0.65	11.6	0.66	39.3	
4	0.67	88.8	38.6	0.66	39.8	0.64	77.3	
5	0.65	75.5	34.1	0.67	35.0	0.67	50.3	
Mean	0.66	77.5	31.2	0.66	31.6	0.66	67.5	

( $\pm$ SD)	(0.01)	(11.1)	(14.2)	(0.01)	(13.1)	(0.01)	(22.0)
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Values are means for each subject of all the maneuvers performed. Ti/Ttot, duty cycle;  $P_{di}$ , transdiaphragmatic pressure;  $V_T$ , tidal volume; IC, inspiratory capacity. All subjects were able to maintain the imposed duty cycle.

- Figure 3 are examples of bar graphs displaying drops in center frequency
- 5      $CF_{di}$ , targeted levels of transdiaphragmatic pressure  $P_{di}$ , diaphragm pressure-time product  $PTP_{di}$  and the associated diaphragm's electrical activity  $EAdi$  observed during the three volume and lower-pressure expulsive and higher pressure expulsive maneuvers performed in the study. The bars of the graphs of Figure 3 are average values obtained for the five subjects ( $\pm$  SD (Standard
- 10    Deviation)).

As shown in Table 1 and Figure 3, subjects were able to achieve and maintain similar target levels of transdiaphragmatic pressure  $P_{di}$  during the volume maneuver (high lung volume) and the lower-pressure expulsive maneuver at EELV. During the volume maneuver, subjects inspired to an average of  $77.5 \pm 11.1\%$  of their inspiratory capacity. In order to generate the same diaphragm pressure-time product  $PTP_{di}$  at different lung volumes, the volume maneuver (neuromechanical uncoupling) required a diaphragm's electrical activity  $EAdi$  of  $60 \pm 8\%$  of maximum compared to  $25 \pm 8\%$  for the expulsive lower-pressure maneuver at EELV. As shown in Table 2 and Figure 3, despite a matching of the diaphragm pressure-time product  $PTP_{di}$ , the volume maneuver promoted a 17% larger drop in the center frequency  $CF_{di}$  than the expulsive low-pressure maneuver at EELV. Figure 4 are examples of graphs from one representative subject showing the center frequency  $CF_{di}$ , the root-means-square (RMS) of the diaphragm's electrical activity  $EAdi$  and the diaphragm pressure-time product  $PTP_{di}$  plotted over time during the volume maneuver (circles) and the two expulsive maneuvers at end-expiratory lung volume (EELV), one targeting a lower transdiaphragmatic pressure  $P_{di}$  (squares) and the other a higher  $P_{di}$  transdiaphragmatic pressure  $P_{di}$  (triangles). Figure 4 shows that, for the representative subject, the center frequency  $CF_{di}$  declines more rapidly and to a greater extent during the volume maneuver (circles), which

required a high diaphragm's electrical activity EAdi for a similar diaphragm pressure-time product PTP<sub>di</sub>, compared to the expulsive lower-pressure maneuver (squares).

5

Table 2

*Individual CF<sub>di</sub> values observed at end of each maneuver*

Subject	Volume maneuver	Expulsive maneuver Low pressure	Expulsive maneuver High pressure
1	68.3 ± 8.7	81.3 ± 7.9	61.6 ± 5.6
2	67.4 ± 4.5	86.7 ± 8.0	70.5 ± 13.2
3	82.0 ± 12.3	94.3 ± 7.8	71.5 ± 5.0
4	72.8 ± 5.8	89.4 ± 6.1	73.4 ± 3.1
5	80.4 ± 9.3	104.1 ± 4.4	83.6 ± 5.4
<b>Mean±SD</b>	<b>74.2±6.8</b>	<b>91.2±8.6</b>	<b>72.1±7.9</b>

Values are means for each subject for each of the maneuvers performed.

In order to produce a similar drop in center frequency CF<sub>di</sub> during the expulsive maneuver at EELV as was observed during the volume maneuver,  
10 more than a two-fold increase in the target transdiaphragmatic pressure P<sub>di</sub> was required. This was associated with an increase in diaphragm's electrical activity EAdi from 25±8 % to 44±9% of maximum. As can be seen in Figure 4, the rate of decline of the center frequency CF<sub>di</sub> was similar for the volume maneuver (circles) and the expulsive higher-pressure maneuvers (triangles).

15

Presented in Table 3 are the values of transdiaphragmatic pressure P<sub>di</sub>, diaphragm's electrical activity EAdi and center frequency CF<sub>di</sub> for the test-retest of the volume maneuver. During the retest, subjects successfully targeted a transdiaphragmatic pressure P<sub>di</sub> that was similar to that generated during the initial volume maneuver (ICC=0.95). The diaphragm's electrical activity EAdi was also similar (ICC=0.93) as was the drop in center frequency CF<sub>di</sub> (ICC=0.98).

Table 3

5 *Test-retest of the volume maneuver*

	CFdi <sub>o</sub>			CFdi			EAdi			Pdi		
	(Hz)			(Hz)			(% max)			(cm H <sub>2</sub> O)		
Sub- ject	Vol 1		Vol 2	Vol 1		Vol 2	Vol 1		Vol 2	Vol 1		Vol 2
1	90.4		93.1	68.3		66.2	46.5		38.1	47.8		44.6
2	102.5		108.2	67.4		65.7	42.4		44.4	38.6		38.3
3	94.7		100.3	82.0		84.7	49.8		55.0	34.1		30.7
4	99.7		99.5	72.6		72.1	51.3		50.6	36.5		35.8
Mean	96.8		100.3	72.8		71.9	66.7		65.0	25.3		29.5
(± SD)	(5.4)		(6.2)	(6.7)		(8.8)	(10.7)		(11.9)	(9.4)		(7.0)
ICC		0.94			0.98			0.93			0.95	

EAdi, diaphragm electrical activity calculated as root-mean-square; CFdi<sub>o</sub>, baseline center frequency determined during resting breathing; CFdi plateau value of the center frequency at the end of the volume maneuver; P<sub>di</sub>, transdiaphragmatic pressure; Vol 1, first volume maneuver performed; Vol 2, second volume maneuver performed; ICC, interclass correlation coefficient.

10 coefficient.

## DISCUSSION

The study evaluated intermittent static contractions maintained at two different lung volumes, in order to examine the effect of altered neuromechanical coupling and increased diaphragm electrical activation, on diaphragm sarcolemma excitability, assessed by changes in center frequency CF<sub>di</sub>. It was

found that, for a given targeted diaphragm pressure-time product  $PTP_{di}$ , the drop in center frequency  $CF_{di}$  was greater when the diaphragm's electrical activity  $EAdi$  was increased by neuromechanical uncoupling, suggesting that the level of muscle activation influences the center frequency  $CF_{di}$ .

5

Studies on the canine diaphragm have demonstrated that changes in center frequency  $CF_{di}$  are associated with changes in the mean action potential conduction velocity (APCV) [38], confirming previous mathematical models [29]. During muscle contractions, both center frequency  $CF_{di}$  and muscle fiber APCV depend to a smaller extent on the cable properties of the fiber [38, 39], and to a larger extent on the muscle membrane excitability [17, 18, 29, 39]. The excitability of the muscle fiber membrane is dependent on the trans-membrane gradient of potassium, and with increased muscle activation, efflux of potassium increases. In order to defend the extra-cellular potassium concentration and hence, the membrane potential, the cell depends on the re-uptake of potassium, e.g. via the ATP (Adenosine TriPhosphate) dependent sodium/potassium pump [12], and washout via the blood circulation [25], i.e. diffusion of potassium from the extra-cellular space into the blood stream. Regardless if blood flow is reduced [23, 31, 42], or the muscle activation is increased, as in the present work, the muscles' electrical activity will indicate reduced membrane excitability, by shifts in the power spectrum toward lower frequencies. The center frequency  $CF_{di}$  can also be affected by factors such as motor unit territory, number of fibers in the motor unit, dispersion in arrival times of the single contributions in the motor unit signal, dispersion in action potential conduction velocities between motor units that can cause the diaphragm's electrical activity  $EAdi$  power spectrum to shift [4, 29]. However, given that these influences are minor in healthy muscles [30] and given that the test situation did not allow for much variability in the contractile pattern, it is unlikely that these influences had more than a minor impact on the results.

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In the study, a constant transdiaphragmatic pressure  $P_{di}$  was targeted with a constant duty cycle at two different lung volumes, and it was therefore

assumed that transdiaphragmatic pressure  $P_{di}$  hindrance to blood flow under those conditions remained relatively similar at the different muscle lengths [19]. However, in order to achieve the same target transdiaphragmatic pressure  $P_{di}$  at an increased lung volume, diaphragm's electrical activity EAdi was increased,  
5 which represents an increase in energy demand/consumption as well as increased metabolic output (e.g. potassium efflux) from the cell. As can be seen in Figure 3, the rate of decline of  $CF_{di}$  at increased lung volume was significantly higher than that observed when the same pressure was targeted at FRC (doubling of transdiaphragmatic pressure  $P_{di}$  at the same lung volume) with  
10 lower diaphragm's electrical activity EAdi. Vitro studies have also demonstrated that increased activation (i.e. demand), accomplished by increasing stimulation frequency of a muscle shortened to 70% of its optimum length, in order to obtain the same tension generated at optimum length, resulted in an increased fatigue in the shortened muscle [14]. The current study therefore demonstrates that the  
15 higher diaphragm activation required for generating the target transdiaphragmatic pressure  $P_{di}$  at an increased lung volume (neuromechanical uncoupling) influences the rate/extent to which center frequency  $CF_{di}$  decays. Further theoretical evidence for the impact of neuromechanical uncoupling on the center frequency  $CF_{di}$  is provided in the following description.

20

In the absence of neuromechanical uncoupling, an increase in transdiaphragmatic pressure  $P_{di}$  is always associated with an increase in diaphragm's electrical activity EAdi. In the above reported study, doubling of transdiaphragmatic pressure  $P_{di}$  at the same lung volume (FRC) was associated  
25 with an increase in diaphragm's electrical activity EAdi from 25% to 44% of the maximum. Beck et al [6] showed that diaphragm's electrical activity EAdi in absolute values is closely related to transdiaphragmatic pressure  $P_{di}$ , such that activation increases (i.e. energy demand increases) when pressure increases (i.e. energy supply decreases). However, this relationship is altered when the  
30 muscle length changes. In such a circumstance, the transdiaphragmatic pressure  $P_{di}$  continues to reflect diaphragm's electrical activity EAdi only when the transdiaphragmatic pressure  $P_{di}$  is normalized to the maximum

transdiaphragmatic pressure  $P_{di}$  obtained at each corresponding lung volume [6]. It was previously shown that when the same diaphragm's electrical activity  $EAdi$  is targeted at different lung volumes, the higher resulting transdiaphragmatic pressure  $P_{di}$  generated at FRC promotes a greater drop in 5 center frequency  $CF_{di}$  than does the lower pressure produced at the higher lung volume [42]. Such results indicate that for a given neural activation, an increase in force or transdiaphragmatic pressure  $P_{di}$  reduces diaphragm excitability. Consequently, the use of the  $TT_{di}$  and pressure-time product as indices for predicting changes in the excitability of the diaphragm sarcolemma (as reflected 10 by center frequency  $CF_{di}$ ) is limited to conditions of constant neuromechanical coupling, where the diaphragm force generating capacity remains unaltered.

Consistent with previous studies [3, 16, 21, 28], doubling of the target transdiaphragmatic pressure  $P_{di}$  at FRC in the present study increased the rate 15 of decline of the center frequency  $CF_{di}$  as well as the level to which it declined (Figures 3 and 4). This is partially explained by the increase in diaphragm's electrical activity  $EAdi$ , as discussed above. However, it is also partially explained by the fact that:

- 20        i)      diaphragm contractions with a higher transdiaphragmatic pressure  $P_{di}$  tend to hinder blood flow (i.e. energy supply) relatively more than contractions producing a lower transdiaphragmatic pressure  $P_{di}$  [19]; and
- 25        ii)     impaired blood flow to a muscle has the propensity to promote shifts in the electromyographic power spectrum toward lower frequencies [22, 30].

*Methodological and technical aspects*

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In the study the contraction and relaxation periods were maintained at a fixed duration and therefore any potential influence of duty cycle on muscle

function [2, 22] was controlled for. It must be emphasized that accurate physiological measurement of the center frequency  $CF_{di}$  depends on being able to control for:

- 5       (a) changes in muscle-to-electrode distance;
- (b) electrode positioning with respect to the muscle fiber direction and location;
- 10      (c) electrode configuration;
- (d) signal to noise ratio;
- 15      (e) influence of cross-talk from other muscles (including the heart and the esophagus); and
- (f) electrode movement-induced artifacts [7, 36, 38, 39, 40].

In the study, the technology used to measure the power spectrum of the diaphragm's electrical activity EAdi spectrum included means for minimizing these influences [1, 36, 40]. The findings that evoked muscle action potentials are influenced by changes in lung volume [5, 15] have contributed to the assumption of a potential-inherent inaccuracy of measured amplitudes of the diaphragm's electrical activity EAdi [5, 15] and the center frequency  $CF_{di}$  [5].  
20  
25 However, during mild voluntary muscle contractions that do not alter diaphragm membrane excitability, it has been shown that chest wall configuration/lung volume and changes in muscle length have no effect on diaphragm's electrical activity EAdi and center frequency  $CF_{di}$  [5, 6, 7, 17, 39]. Therefore the above-discussed effect of chest wall configuration/lung volume likely did not have an impact on the results.  
30

Another factor that could have influenced the results of the study is the

difference in partitioning the esophageal and gastric pressures for the same transdiaphragmatic pressure  $P_{di}$  during the various maneuvers. In a previous study [42], where subjects targeted the same diaphragm's electrical activity EAdi at higher and lower lung volumes, greater decreases in center frequency  $CF_{di}$

5 were consistently observed at EELV (higher transdiaphragmatic pressure  $P_{di}$ ), regardless of whether subjects performed an expulsive (i.e. transdiaphragmatic pressure  $P_{di}$  generated mainly by gastric pressure) or a Mueller maneuver (i.e. transdiaphragmatic pressure  $P_{di}$  generated mainly by esophageal pressure) at EELV [42]. In a pilot trial to that study (unpublished observations), it was found

10 that diaphragm contractions generating identical transdiaphragmatic pressure  $P_{di}$ , duty cycle and diaphragm's electrical activity EAdi, produced the same trajectory of decrease in center frequency  $CF_{di}$ , whether subjects performed expulsive or Mueller maneuvers. Therefore, it is not believed that differences in the partitioning of the esophageal and gastric pressures during the volume and

15 expulsive maneuvers in the current study had an effect on the outcomes observed.

#### *Clinical implications*

20 The results of the above reported study have direct implications to subjects or patients being weaned from mechanical ventilation. It is well known that patients undergoing a weaning trial may demonstrate dynamic changes in EELV (dynamic hyperinflation) [43], which similar to the study would alter the neuromechanical coupling of the diaphragm. In order to compensate for this

25 uncoupling (i.e. maintain the same transdiaphragmatic pressure  $P_{di}$ ), the patient would need to increase diaphragm activation. The combination of an increased activation of the patient's diaphragm, with an elevated transdiaphragmatic pressure  $P_{di}$  would, according to the present study, lead to decreased center frequency  $CF_{di}$  (excitability), and possibly an increased respiratory effort

30 sensation [42]. Shifts in the H/L ratio of the power spectrum of the diaphragm's electrical activity EAdi have been reported in patients with respiratory failure in whom ventilatory assistance is removed [8, 13]. However, given that diaphragm

weakness is prevalent in mechanically ventilated patients [24], it remains to be determined what combined levels of diaphragm's electrical activity EAdi and transdiaphragmatic pressure  $P_{di}$  would affect center frequency  $CF_{di}$ .

5

## CONCLUSION

The above-reported study shows that diaphragm activation can be used to determine diaphragm membrane excitability and changes in center frequency  $CF_{di}$ . Furthermore it shows that the diaphragm pressure-time product  $PTP_{di}$  and 10 tension-time index  $TT_{di}$  cannot be considered as valid reflections of diaphragm energy consumption and/or sarcolemma excitability when neuromechanical coupling is altered.

With data from the above investigation or study, the diaphragmatic 15 muscle force can be estimated from measurements of the diaphragm's electrical activity EAdi in two ways.

A first way for estimating the diaphragmatic muscle force uses the following equation:

20

$$F = \mu EAdi \quad (1)$$

where F is the diaphragmatic muscle force,  $\mu$  is a proportionality constant, and 25 EAdi is a measure of the signal strength of the electrical activity of the patient's diaphragm. Here the square root of the first power spectral moment is used since it represents the signal strength, which has been compensated for the influence of changes in the propagation velocity of the myoelectric action potentials [29].

30

A second way for estimating the diaphragmatic muscle force uses the spectral changes during diaphragm contraction. For a forceful periodic muscle loading, the center frequency  $CF_{di}$  decreases from an initial center frequency

$CF_0$  to a final plateau value  $CF_\infty$  according to the equation [26]:

$$CF_\infty = CF_o (1 - \kappa) T_D / [(1 - \kappa) T_D + \kappa T_R] \quad (2)$$

- 5 where  $\kappa$  is the duty cycle, i.e. the inspiration time in relation to the total time period, and  $T_R$  is the center frequency  $CF_{di}$  recovery time constant pertaining to an approximately exponential time curve which is rather independent of the muscle force [9]. The symbol  $T_D$  denotes the time constant for the decrease in center frequency  $CF_{di}$ , which is related to the muscle force as [27]:

10

$$T_D = \eta / (F - F_c) \quad (3)$$

- In this equation  $\eta$  is a proportionality constant and  $F_c$  is a critical force level above which muscle fatigue starts to develop. Equation (2) is rearranged to 15 obtain the experimentally determinable quantity:

$$Q = T_R / T_D = [(1 - \kappa) / \kappa] [(CF_0 - CF_\infty) / CF_\infty] \quad (4)$$

Equations (3) and (4) then give:

20

$$F = F_c + Q\eta / T_R \quad (5)$$

Making equal the two force estimates of equations (1) and (5) the following relation is found:

25

$$\alpha EAdi - \beta - Q = 0 \quad (6)$$

where

30

$$\alpha = \mu T_R / \eta \quad (7)$$

and

$$\beta = F_C T_R / \eta \quad (8)$$

5

Relation (6) represents a set of three equations (for the three experimental conditions) with two unknowns. A fitting procedure with data from the following Table 4 with simultaneous minimization of the relative errors in the diaphragm's electrical activity signal strength EAdi and the quantity Q, gives the values  $\alpha = 10$  0.00417 and  $\beta = 0.0419$  with a relative fitting error of 0.24.

Table 4  
Experimental results and calculated values

	$P_{di}$ (cm H <sub>2</sub> O)	EAdi (a.u.)	CF <sub>0</sub> (Hz)	CF <sub>∞</sub> (Hz)	Q	Force ratios			Geometric factors		
						$\phi_I$	$\phi_{II}$	$\phi_m$	$\gamma_I$ (cm H <sub>2</sub> O)	$\gamma_{II}$ (cm H <sub>2</sub> O)	$\gamma_m$
Volume Maneuver	31.2	60.9	100	74.2	0.175	6.12	5.20	5.66	122	144	133
Expulsive maneuver (Lower pressure)	31.6	24.9	100	91.2	0.050	2.50	2.19	2.35	303	347	325
Expulsive maneuver (Higher pressure)	67.5	44.3	100	72.1	0.196	4.45	5.69	5.07	364	285	324

15  $P_{di}$ , transdiaphragmatic pressure; a.u., arbitrary units; EAdi, signal strength of the electrical activity of the diaphragm; CF<sub>di0</sub>, baseline diaphragm center frequency determined during resting breathing; CF<sub>di</sub>, plateau value of the diaphragm center frequency at the end of the maneuver; Q, ratio of the time constants of CF<sub>di</sub> recovery and decline, see equation (5);  $\phi_I$ , see equation (9);  $\phi_{II}$ , see equation (10);  $\phi_m$ , mean of  $\phi_I$  and  $\phi_{II}$ ;  $\gamma_I$ , see equation (14);  $\gamma_{II}$ , see equation (15);  $\gamma_m$ , mean of  $\gamma_I$  and  $\gamma_{II}$ .

20

With  $\alpha$  and  $\beta$  known, the experimental values of the diaphragmatic muscle force F can be expressed in relation to the critical force level  $F_C$  for onset

of deterioration of cell excitability, i.e. the critical force level above which muscle fatigue starts to develop. The two ways to describe this are obtained by rearranging equations (1) and (7), and equation (5), respectively, which gives:

$$5 \quad \phi_I = (F / F_c)_I = \alpha EAdi / \beta \quad (9)$$

and

$$10 \quad \phi_{II} = (F / F_c)_{II} = 1 + Q / \beta \quad (10)$$

These quantities have been determined and are listed in Table 4 together with their mean values  $\phi_m$ .

15 The observed transdiaphragmatic pressure  $P_{di}$  is assumed to be related to the diaphragmatic muscle force  $F$  as:

$$P_{di} = F G \quad (11)$$

20 where  $G$  is a geometrical factor taking into account that the diaphragm muscle changes its shape with the inspired volume. This factor  $G$  is thus assumed to be the same during the expulsive maneuvers with lower or higher  $P_{di}$  production performed at end-expiratory lung volume. As with the force relations, the transdiaphragmatic pressure  $P_{di}$  can be expressed in two ways, relating to the diaphragm's electrical activity signal strength  $EAdi$  and to the fatigue induced 25 spectral changes. Combining equations (1), (5), and (11) leads to the following relations:

$$P_{di} = \mu EAdi G \quad (12)$$

and

30

$$P_{di} = (F_c + Q\eta / T_R)G \quad (13)$$

Relations (12) and (13) can be further developed with relations (7) and (8) into the two following relations:

$$5 \quad \gamma_I = (G\eta/T_R)_I = Pdi/(\alpha Eadi) \quad (14)$$

and

$$\gamma_{II} = (G\eta/T_R)_{II} = Pdi/(\beta + Q) \quad (15)$$

- 10 Numerical values, calculated for the two expressions, are given in Table 4  
together with their mean values  $\gamma_m$ .

From the results listed in Table 4, it can be concluded that the diaphragmatic muscle force F in relation to the critical force level  $F_c$  are  
15 approximately the same during the volume maneuver and the higher pressure  
expulsive maneuver, which is also reflected in their deterioration of cell  
excitability, expressed by the factor Q. During all conditions the diaphragmatic  
muscle forces F are above the critical force level  $F_c$  as shown by values of  $\phi_m$  in  
Table 4. The geometrical dependence, expressed by the factor  $\gamma_m$ , is obviously  
20 the same during lower pressure expulsive maneuver and higher pressure  
expulsive maneuver, but is much less during the volume maneuver. The ratio  
between the  $\gamma$  values in the volume maneuver and the expulsive maneuvers is  
about 0.41. Since the  $\eta$  values and the  $T_R$  values are expected to be  
independent of the maneuvers, this means that also the factors G have the  
25 same ratio. This indicates a much lower efficiency to convert force into pressure  
during the volume maneuver. The tension time index  $TT_{di}$ , taking into account  
the timing and the pressure, is thus not sufficient to describe the complexity of  
the fatigue development. At least it has to be modified with a volume dependent  
correction factor. Better, though, are methods reflecting the deterioration of cell  
30 excitability and not the mechanical result of the contraction.

*Electromyographic and mechanical methods to detect muscle fatigue*

Based on the above results, techniques to determine critical levels of muscle fatigue during periodic loading (such as respiration) will be described. A 5 number of equations relating certain physiological variables to each other are needed and they will be derived prior to the description of these techniques.

*Periodic muscle load characteristics*

10 Consider a periodic muscle loading, such as the respiratory work, in which repeated muscle contractions alternate with muscle relaxations. The periodic muscle loading is characterized by a time period  $T_0$  and its two parts: the duration of muscle contraction  $T_1$  and the duration of muscle relaxation  $T_2$  where:

15

$$T_0 = T_1 + T_2 \quad (16)$$

In order to simplify the equations, the duty cycle  $\kappa$  is determined as:

20

$$\kappa = T_1 / T_0 \quad (17)$$

The mean diaphragmatic muscle force developed during the time interval  $T_1$  is denoted  $F$ .

25

*Myoelectric changes due to fatigue*

Isometric fatiguing contractions cause the center frequency  $CF_{di}$  of the diaphragm's electrical activity  $EAdi$  diaphragm's electrical activity to decrease exponentially from its resting value  $CF_0$  with a time constant  $T_F$ . During recovery 30 the center frequency  $CF_{di}$  returns gradually to its normal value following an approximately exponential course, described by the recovery time constant  $T_R$ . It is observed that many other characteristics of the power spectrum of the

diaphragm's electrical activity EAdi exhibit the same dependencies such as the median frequency, the zero crossing density, the so-called hi-over-low value, etc. The recovery time constant depends mostly on the density of capillaries in the muscle and is rather insensitive to the exerted force. The fatigue time 5 constant is strongly dependent on the force when it exceeds a certain critical level  $F_c$ . The relation is:

$$T_F = \eta / (F - F_c) \quad \text{for } F > F_c \quad (18a)$$

10 and

$$T_F \rightarrow \infty \quad \text{for } F \leq F_c \quad (18b)$$

The combination of repeated work and recovery events causes the center 15 frequency CFdi to decrease from the initial value to a final plateau value  $CF_\infty$ , at which there is a balance between the metabolite production during work and wash-out during recovery. The plateau value is:

$$CF_\infty = CF_0 (1 - \kappa) T_F / [(1 - \kappa) T_F + \kappa T_R] \quad (19)$$

20

Introducing the notations:

$$\Delta CF = CF_0 - CF_\infty \quad (20)$$

25 and

$$\varepsilon = \Delta CF / CF_0 \quad (21)$$

Equation (19) can then be rearranged to read:

30

$$\kappa = 1 / [1 + (T_R / T_F)(\Delta CF / CF_\infty)] \quad (22)$$

With the notation:

$$Q = T_R / T_F \quad (23)$$

5 it is found that:

$$Q = [(1 - \kappa) / \kappa] \Delta CF / CF_\infty \quad (24)$$

which is an experimentally measurable quantity.

10

#### *Force and pressure*

The diaphragmatic muscle force  $F$  can be determined for skeletal muscles working over joints without synergistic effects from other muscles. For 15 the diaphragm muscle the force cannot be directly measured, rather the transdiaphragmatic pressure  $P_{di}$  is obtained as a proportional measure. The following relation could be used:

$$F = \mu E \quad (25)$$

20

where  $\mu$  is a proportionality constant and  $E$  is the signal strength of the diaphragm's electrical activity  $E_{Adi}$ , preferably based on the first spectral moment which is rather insensitive to metabolic changes caused by fatigue. The relation to the pressure is proportional but non-linear. This fact is taken into 25 consideration by introducing the factor  $G(V)$  which is volume ( $V$ ) dependent, i.e.:

$$P_{di} = F G(V) \quad (26)$$

Thus,

30

$$\mu G(V) = P_{di} / E \quad (27)$$

which also is an experimentally measurable quantity.

*Myoelectric signal strength and spectral changes*

5 Rearrangement of equation (18a) and insertion of equations (23) and (25)  
gives:

$$\alpha E - \beta - Q = 0 \quad (28)$$

10 where

$$\alpha = \mu T_R / \eta \quad (29)$$

and

15

$$\beta = F_c T_R / \eta \quad (30)$$

It can be observed that  $\alpha$  is dependent, through the parameter  $\mu$ , on the  
electrode geometry and placement in relation to the muscle, while the other  
20 parameters are rather constant for similar muscles.

Experiments under fatiguing conditions at any volume give corresponding  
values of  $E$  and  $Q$  (through the center frequency changes). A data fitting  
procedure (not regression) gives numerical values to  $\alpha$  and  $\beta$ . With  $\alpha$  and  $\beta$   
25 known, an estimate of the diaphragmatic muscle force  $F$  can be obtained in  
relation to its fatigue threshold value, i.e.:

$$F/F_c = E \alpha / \beta \quad (31)$$

30 As long as  $F/F_c$  is smaller than one, isometric fatigue of the patient's  
muscle does not develop. That means that the signal strength should be lower  
than the critical value:

$$E < E_{ISOM} = \beta / \alpha \quad (32)$$

For periodic muscle work, higher forces and signal levels are tolerable.

5

*Spectral changes as indicators of tolerable concentration of metabolites*

The relative spectral change  $\varepsilon$  of the diaphragm's electrical activity EAdi, defined in equation (21), is an indirect measurement of remaining concentration 10 of metabolites in the muscle during periodic fatiguing contractions. It seems that the muscle very rapidly goes into an anaerobic metabolic state once the force is higher than  $F_c$  and that virtually all contractions above this level causes changes in the center frequency CFdi. Therefore it is likely that a certain small value of  $\varepsilon$  is tolerable as long as it is below a certain critical level, which we denote  $\varepsilon_c$ . 15 With this critical value introduced into equation (22) and simultaneous use of equations (18a) and (23), it can be found that a condition for long term fatigue not to occur is:

$$\kappa < 1 / \{1 + [(1 - \varepsilon_c) / \varepsilon_c] T_R (F - F_c) / \eta\} \quad (33)$$

20

This expression can be rearranged to give the force condition:

$$F < F_c + [(1 - \kappa) / \kappa] [\varepsilon_c / (1 - \varepsilon_c)] \eta / T_R \quad (34)$$

25 or, together with equation (30),

$$F < F_c \{1 + [(1 - \kappa) / \kappa] [\varepsilon_c / (1 - \varepsilon_c)] / \beta\} \quad (35)$$

Since the force in diaphragmatic contractions cannot be simply measured, 30 equations (33) to (35) are expressed as functions of the signal strength E and the transdiaphragmatic pressure Pdi. Use of equations (25) and (26) give for the signal strength E of the diaphragm's electrical activity EAdi:

$$\kappa < 1 / \{1 + [(1 - \varepsilon_c) / \varepsilon_c] (\alpha E - \beta)\} \quad (36)$$

and

5

$$E < \{\beta + [(1 - \kappa) / \kappa] [\varepsilon_c / (1 - \varepsilon_c)]\} / \alpha \quad (37)$$

and for the transdiaphragmatic pressure Pdi:

10

$$\kappa < 1 / \{1 + [(1 - \varepsilon_c) / \varepsilon_c] (\alpha P_{di} - \beta)\} \quad (38)$$

and

15

$$P_{di} < \mu G(V) \{\beta + [(1 - \kappa) / \kappa] [\varepsilon_c / (1 - \varepsilon_c)]\} \quad (39)$$

*NON-RESTRICTIVE ILLUSTRATIVE EMBODIMENT OF A METHOD  
AND DEVICE FOR DETERMINING AN OPTIMAL LEVEL OF  
VENTILATORY ASSIST TO A VENTILATOR DEPENDENT PATIENT*

20

Non-restrictive illustrative embodiments of the method and device for determining an optimal level of ventilatory assist to a ventilator-dependent patient will now be described.

25 First embodiment of Figure 5

*Operation 501*

The signal strength of the diaphragm's electrical activity EAdi is 30 monitored through a detector 502. As illustrated in Figure 1, detector 502 may comprise, for example, a computer 4 to measure the signal strength of the electrical activity EAdi of the patient's diaphragm through a linear array 5 of

electrodes mounted on an esophageal catheter 6 inserted through the patient's nostril (or patient's mouth) until the electrode array 5 is positioned in the gastro-esophageal junction 10 of the patient's diaphragm 7.

5

*Operation 502*

A calculator 503 calculates the coefficients  $\alpha$  and  $\beta$  using equation (28):

$$10 \quad \alpha E - \beta - Q = 0 \quad (28)$$

with myoelectric data from fatigue tests (calibration). Fatigue test can be performed by either reducing the level of assist, or performing a short airway occlusion, while measuring the myoelectric activity during a few inspiratory attempts. To shorten and facilitate the fatigue test the subject could be encouraged to voluntarily increase his efforts. Such a test is routinely performed to determine the maximum inspiratory airway pressure.

## *Operation 504*

20

The calculator 503 calculates the duty cycle  $\kappa$  as described hereinabove.

## *Operation 505*

25 The calculator-503 calculates estimates of a critical level of the relative  
 spectral change  $\varepsilon_c$  of the diaphragm's electrical activity  $EAd_i$  from the general  
 experimental fact that fatigue does not occur below a duty cycle of 0.2 even at  
 maximum muscle force and that the critical force level  $F_c$  is approximately 0.2  
 times the maximum force. Equation (33) then gives  $\varepsilon_c \approx \beta/(\beta+8/9)$ , or, since both  
 30  $\varepsilon_c$  and  $\beta$  are small quantities:

$$\varepsilon_C \approx \beta \quad (40)$$

*Operation 506*

The calculator 503 calculates a critical signal strength of the diaphragm's  
5 electrical activity EAdi above which isometric muscle fatigue develops, using the  
relation:

$$E < E_{ISOM} = \beta / \alpha \quad (32)$$

10

*Operation 507*

If myoelectric monitoring is used (giving signal strength and duty cycle),  
the calculator 503 calculates a critical signal strength of the diaphragm's  
15 electrical activity EAdi above which long term muscle fatigue develops, is  
calculated using equation (37):

$$E < \{ \beta + [(1 - \kappa) / \kappa] [\varepsilon_c / (1 - \varepsilon_c)] \} / \alpha \quad (37)$$

20

*Operation 508*

A controller 509 controls the ventilatory assist, for example the gain of the  
ventilatory assist at a level such that the signal strength of the diaphragm's  
electrical activity EAdi does not exceed that described in relation (37) (higher  
25 support suggest unnecessary muscle inactivation) to prevent long-term muscle  
fatigue to develop::

$$E < \{ \beta + [(1 - \kappa) / \kappa] [\varepsilon_c / (1 - \varepsilon_c)] \} / \alpha \quad (37)$$

30

However, the signal strength of the diaphragm's electrical activity EAdi  
should not exceed that described in equation (32) (this level indicates the level  
for muscle fatigue during isometric contractions) to prevent isometric muscle

fatigue to develop::

$$E < E_{ISOM} = \beta/\alpha \quad (32)$$

5 Second embodiment of Figure 6

*Operation 601*

The signal strength of the diaphragm's electrical activity EAdi is  
10 monitored through a detector 602. As illustrated in Figure 1, detector 602 may comprise, for example, a computer 4 to measure the signal strength of the electrical activity EAdi of the patient's diaphragm through a linear array 5 of electrodes mounted on an esophageal catheter 6 inserted through the patient's nostril (or patient's mouth) until the electrode array 5 is positioned in the gastro-  
15 esophageal junction 10 of the patient's diaphragm 7.

*Operation 603*

A detector 604 monitors the patient's transdiaphragmatic pressure P<sub>di</sub>. As  
20 illustrated in Figure 1, detector 604 may comprise, for example, a computer 4 to continuously measure the transdiaphragmatic pressure P<sub>di</sub> by detecting the esophageal P<sub>es</sub> and gastric P<sub>ga</sub> pressures through respective gastric 8 and esophageal 9 balloons mounted on the catheter 6 on opposite sides of the array 5 of electrodes, and by processing the detected esophageal P<sub>es</sub> and gastric P<sub>ga</sub> pressures to obtain the patient's transdiaphragmatic pressure P<sub>di</sub>.  
25

*Operation 605*

A calculator 606 calculates the coefficients  $\alpha$  and  $\beta$  using equation (28):  
30

$$\alpha E - \beta - Q = 0 \quad (28)$$

with myoelectric data from fatigue tests (calibration). Fatigue test can be performed by either reducing the level of assist, or performing a short airway occlusion, while measuring the myoelectric activity during a few inspiratory attempts. To shorten and facilitate the fatigue test the subject could be  
5 encouraged to voluntarily increase his efforts. Such a test is routinely performed to determine the maximum inspiratory airway pressure.

*Operation 607*

10 The calculator 606 calculates the duty cycle  $\kappa$  as described hereinabove.

*Operation 608*

The calculator 606 calculates estimates of a critical level of the relative  
15 spectral change  $\varepsilon_c$  of the diaphragm's electrical activity EAdi from the general experimental fact that fatigue does not occur below a duty cycle of 0.2 even at maximum muscle force and that the critical force level  $F_c$  is approximately 0.2 times the maximum force. Equation (33) then gives  $\varepsilon_c \approx \beta/(\beta+8/9)$ , or, since both  $\varepsilon_c$  and  $\beta$  are small quantities:  
20

$$\varepsilon_c \approx \beta \quad (40)$$

*Operation 609*

25 The calculator 606 calculates a critical signal strength of the diaphragm's electrical activity EAdi above which isometric muscle fatigue develops, using the relation:

$$E < E_{ISOM} = \beta/\alpha \quad (32)$$

30

*Operation 610*

The calculator 606 calculates a critical level of the transdiaphragmatic pressure  $P_{di}$  using relation (39):

$$P_{di} < \mu G(V) \{ \beta + [(1 - \kappa) / \kappa] [\varepsilon_c / (1 - \varepsilon_c)] \} \quad (39)$$

5

Operation 610 requires knowledge about the geometrical  $G(V)$  dependence. This factor  $G(V)$  can be obtained from a calibration of the experimentally measurable quantity  $P_{di} / E$  as shown in equation (27):

$$10 \quad \mu G(V) = P_{di} / E \quad (27)$$

Alternatively, geometrical dependence  $G(V)$  of inspiratory pressure can also be estimated by performing single or multiple breath airway occlusions at two lung volumes, e.g. end-inspiration and end expiration lung volumes, while 15 the volume difference is measured by a computer with, for example, at least one flow meter (see computer 4 and pneumotachograph 3 of Figure 1).

#### *Operation 611*

20 A controller 612 controls the ventilatory assist, for example the gain of the ventilatory assist at a level such that:

- the monitored signal strength of the diaphragm's electrical activity  $E_{Adi}$  does not exceed that described in relation (32) (this level indicates the level for 25 muscle fatigue during isometric contractions) to prevent isometric muscle fatigue to develop:

$$E < E_{ISOM} = \beta / \alpha \quad (32)$$

30 - the monitored patient's transdiaphragmatic pressure  $P_{di}$  does not exceed that described in relation (39) to prevent long-term muscle fatigue to develop:

$$P_{di} < \mu G(V) \{ \beta + [(1 - \kappa) / \kappa] [\varepsilon_c / (1 - \varepsilon_c)] \} \quad (39)$$

Although the present invention has been described hereinabove with reference to non-restrictive illustrative embodiments thereof, it should be kept in mind that these embodiments can be modified at will within the scope of the appended claims without departing from the spirit and nature of the present invention. In particular but not exclusively:

- the present invention pertains not only to CFdi and RMS but possibly to other types of measures;
- the present invention can be implemented through measurement of the electrical activity of respiration-related muscles other than the diaphragm; and
- the present invention is concerned with any method of mechanical ventilation, including negative pressure ventilation.

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